mechanisms of morgane Reations, a study of vistal corpores in solution. 1967

Hard and Soft Acids and Bases

The results of numerous investigations for the first transition series she that, regardless of the nature of the donor group, the so-called natural order of the stability of complexes of bivalent transition metals is Mn < Fe < Co < Ni < Cu > Zn. Only in a very few cases he deviations from this order been observed. It has been found, for example that the stability of $Fe(phen)_a^{3+}$ is greater than expected and, furthermooghat the value of K_a for the addition of the third group is larger than the of either K_1 or K_2^{98} . This is attributed to an electronic rearrangement, for it was shown that the bis complex, $Fe(phen)_2 X_2$, is paramagnetic, where the tris complex, $Fe(phen)_3^{2+}$, is diamagnetic.

If we attempt to list the other metal ions in the order of increasing stability of their complexes, this turns out to be impossible unless the ligand is specified. In the same way, it is not possible to put the comming and into an order of complexing ability unless a reference metal ion is specified.

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Various metal ions fall into two categories: (1) those binding strong y to bases which bind strongly to the proton, that is, basic in the usu'll sense; (2) those binding strongly to highly polarizable or unsaturated bases, which often have negligible proton basicity. Division into the etwo categories is not absolute and intermediate cases occur, but the classification is reasonably sharp and appears to be quite useful. It is convenient to divide bases also into two categories, those that are polarifable or "soft," and those that are non-polarizable or "hard." It is possible for a base to be both soft and strongly binding toward the proton, e.g. sulfide ion. Still it is generally true that hardness is associated with good proton binding. For example, for the bases in which the coordinating atom is from Groups V, VI, and VII (the great majority of all bases), the state of the most basic to the proton.

It has long been recognized 101 that metal ions can be divided into two classes depending on whether they form their most stable complexed with the first ligand atom of each group, class (a), or whether they form their most stable complexes with the second or a subsequent member queach group, class (b). Thus, for class (a) metal ions the order of stability is as follows:

 $F^- > Cl^- > Br^- > I^ O \gg S > Se > Tc$ $N \gg P > As > Sb > Bi$

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Whereas for class (1) the order of stability is

$$F^- < Cl^- < Bl^- < I^-$$

 $O \ll S \sim Se \sim Te$

$$N \ll P > As > Sb > Bi$$

class (a) acids, how vcr.

Table 1.6 contails a list of all generalized acids for which sufficient information can by found in the literature to enable a choice between

soft bases.

Class (a) metal io is bind best to the least polarizable (hardest) atom of a family, whereas lass (b) metal ions bind best to a more polarizable (softer) atom of the same family. Notice that it is not always the most polarizable (softest) atom which forms the most stable complexes with a class (b) metal ion. The reason for this is that many bases, such as the stibines, are weak t ises toward all metal ions. Their complexes with class (b) metals will usu lly be much more stable than their complexes with

Other Lewis acid can be examined to see whether the stability of their acid-base adducts corresponds to class (a) or class (b) behavior. The acidbase or donor-acceptor 102 adducts which they form may be inorganic or organic molecules, complex ions, or charge-transfer complexes.

class (a) and class (b) to be made. In classifying Lewis acids, the criterion previously used 101 y as followed whenever possible, that is, to compare the stabilities of fluori le versus iodine, oxygen versus sulfur, and nitrogen versus phosphorus type complexes. When such comparisons are not feasible, other crit ria may be used. 100 One is that class (b) acids will complex readily with a variety of soft bases that are of negligible proton basicity. These include CO, olefins, aromatic hydrocarbons, and the like. The common ch racteristics of the two classes of Lewis acids are easily discernible from 1 able 1.6. The features which bring out class (a) behavior are small sii e, high positive oxidation state, and no easily distorted outer electrons. Class (b) behavior is associated with a low or zero oxidation state, with large size, and with easily distorted outer electrons. Both metals and non-metals can be either (a) or (b) type acids, depending or their charge and size. Since the features which promote class (a) behavior are those leading lo low polarizability, and those which create type (b) behavior lead to high polarizability, it is convenient to call class (a) acids "hard" and class (b) acids "soft." We then have the useful generalization that hard acids pi fer to associate with hard bases, and soft acids prefer

It is not implied that complexes of hard acids and soft bases, or vice versa, cannot exist and be quite st ble. For example, CH₂ is a soft base yet compounds such as Mg(CH₂), can easily be made. Nevertheless this compound is thermodynamically unstable to hydrolys! whereas Hg(CH)2 is thermodynamically stable to hydrolysis.

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Hard and Soft Acids and Bases

Polarizability is simply a convenient property to use as a classification. It may well be that other properties which are roughly proportional to polarizability are more responsible for the typical behavior of the toclasses of acids. For example, a low ionization potential is usually linked to a high polarizability, and a high ionization potential to a low polarizability. Hence, ionization potential or the related electronegativity might the important property. Unsaturation, with the possibility of acceptor

Table 1.6 Classification of Lewis acids

| Hard | Soft | |
|----------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------|
| H+, Li+, Na+, K+ | Cu+, Ag+, Au+, Tl+, Hg+ | |
| Be ²⁺ , Mg ³⁺ , Ca ²⁺ , Sr ³⁺ , Mn ³⁺ | Pd ²⁺ , Cd ³⁺ , Pt ²⁺ , Hg ²⁺ , CH ₈ Hg ⁺ , Co(CN) ₅ ²⁻ , Pt ²⁺ , Tc ²⁺ | |
| Al ³⁺ , Sc ³⁺ , Ga ³⁺ , In ⁸⁺ , La ³⁺ | Tl^{3+} , $Tl(CH_3)_3$, BH_3 , $Ga(CH_8)_8$, | Ga() |
| N ⁸⁺ , Gd ³⁺ , Lu ³⁺ | Gala, InCla | |
| Cr ³⁺ , Co ³⁺ , Fe ³⁺ , As ³⁺ , Ce ³⁺ | RS+, RSe+, RTe+ | - H |
| Si ⁴⁺ , Ti ⁴⁺ , Zr ⁴⁺ , Th ⁴⁺ , Pu ⁴⁺ | I+, Br+, HO+, RO+ | |
| UO_2^{2+} , $(CH_8)_9Sn^{2+}$, VO^{2+} , MoO^{8+} | I ₂ , Br ₂ , ICN, etc. | - |
| $BeMe_2$, BF_3 , $B(OR)_3$ | trinitrobenzene, etc. | - |
| Al(Cll ₂) ₃ , AlCl ₂ , AlH ₂ | chloranil, quinones, etc. | |
| RPO ₂ +, ROPO ₂ + | tetracyanoethylene, etc. | - |
| RSO ₂ +, ROSO ₂ +, SO ₈ | O, Cl, Br, I, N | il |
| 1 ⁷⁺ , I ⁶⁺ , Cl ⁷⁺ , Cr ⁶⁺ | M ^o (metal atoms) | - ! |
| RCO+, CO ₂ , NC+ | bulk metals | |
| HX (hydrogen-bonding molecules) | CH ₈ , carbones | il |

Borderline

Fe²⁺, Co²⁺, Ni²⁺, Cu²⁺, Zn²⁺, Pb²⁺, Sn²⁺, Sb⁸⁺, Bi³⁺, Rh³⁺, Ir⁸⁺, B(CH₃)₈, S NO⁺, Ru³⁺, Os³⁺, R₃C¹, C₆H₆⁺, Ga H₃

bonding in the acid-base complex, and ease of reduction, favoring stroig electron transfer to the acid, are also associated with high polarizabilt.

While considerable variation can exist because of the effect of overall charge, the usual stability order overall for class (b) metal ions is $S \sim C > I > Br > Cl > N > O > F$. For class (a) metal ions a stroig inversion of this order occurs, so that often only oxygen and fluoride complexes can be obtained in aqueous solution. 108 It may be noted that the order given above is that of increasing electronegativity.**

** Table 3.4 lists a series of stability constants for various ligands for the typical sift acid CH₈Hg*.

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